Efficacy of Thermophilic Soil-Isolated *Paenibacillus* sp. NBR10 as a Chitinolytic and Biocontrol Bacterium-In vitro Study

Mohamed H. El-Sayed(1,2), Olfat M. Nassar(3), Heba A. Nasr(4), Abd El-Nasser A. Kobisi(4)*

(1)Department of Biology, Faculty of Science and Arts, Northern Border University, Kingdom of Saudi Arabia; (2)Department of Botany and Microbiology, Faculty of Science (Boys), Al-Azhar University, Cairo, Egypt; (3)Department of Home Economics (Nutrition), Faculty of Science and Arts, Northern Border University, Kingdom of Saudi Arabia; (4)Plant Protection Department, Desert Research Center, El-Mataria, Cairo, Egypt.

Among microbial enzymes, chitinases received increased attentions due to their wide applications especially in agriculture sector for biocontrol of fungal phytopathogens. The growing interest in chitinases with thermostability nature, forced many researchers to isolate and characterize novel chitinase producing-bacteria from extreme environments. The present study describe characterization of thermostable chitinase produced by thermophilic soil-isolated bacterial strain NBR10 from Rafha governorate, Saudi Arabia. Chitinase producing NBR10 strain was identified as *Paenibacillus* sp. according to the traditional methods of morphological, biochemical and physiological characteristics coupling with sequencing of 16S rRNA (GenBank accession number KT957624.1). The isolated strain was found to have a potent antifungal against three of the phytopathogenic fungi *Fusarium oxysporum, Alternaria burnsii* and *Rhizoctonia solani* with percent of growth inhibition 52.5, 75.0 and 85.71%, respectively. Chitinase obtained from *Paenibacillus* sp. NBR10 showed activity at ranges of temperature (30–90°C) and pH (4–9), showing optimum activity at 55°C and pH 7.0, respectively. Also, it exhibited high thermostability at higher temperatures, where 90% of its activity is retained at 65°C for 36h. The promising chitinolytic and antifungal activity of locally isolated *Paenibacillus* sp. NBR10, candidate this strain to be used as a potential biological control agent.

**Keywords:** Soil, *Paenibacillus* sp. NBR10, Thermostable chitinase, Antifungal, Biological control.

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**Introduction**

Every year thousands of tons of biowastes are generated globally (Pachapur et al., 2016). Persistence and accumulation of these wastes cause environmental issues (Kim et al., 2007). After cellulose, chitin considers the second abundant polysaccharide found in nature (Garcia-Fraga et al., 2015). Chitin is found in many sources in nature such as fungal cell walls, shells of crustaceans, exoskeleton of arthropods and eggs of nematodes (Brzezinska et al., 2014) and in midgut of many invertebrates (Hagedus et al., 2009).

Chitin has not been applied to large-scale industrial use because of its extreme insolubility (Jeraj et al., 2006). Breaking down of chitin-containing wastes into oligosaccharides has attracted great attention because the useful biological activities of these oligosaccharides such as antifungal, antitumor, and immunity enhancement activities (Dai et al., 2011).

Chitin is a polysaccharide with linear structure formed of N-acetyl-D-glucosamine (NAG) residues linked with β-1-4 bonds (Zarei et al., 2012), and hydrolyzed by diverse chitinolytic enzymes with different modes of hydrolysis. These enzymes can be classified as endochitinases, exochitinases, chitobiases, and β-N-acetylglucosaminidases (Brzezinska et al., 2014).

Chitinases are produced by different
microorganisms such as bacteria, fungi and algae and are involved in various biological processes as defense against pathogens, nutrition and morphogenesis (Yan & Fong, 2015). Commercially chitinolytic enzymes are used to produce NAG and chitooligosaccharides as part of pharmaceutical formulations. Purified chitinase enzyme has been used to generate protoplasts from yeasts and fungi and to deter the growth of pathogenic fungi, treatment of chitinous wastes (Dahiya et al., 2006).

In the recent decades, bacterial chitinases received over attention because of their promising applications, especially in agriculture sector as abiocontrol agent against harmful phytopathogenic fungi and insects, because of their containing of chitin components. Moreover, chitinases form bacteriaact as a potent alternative biopesticides for the chemical fungicides (Singh, 2010 and Yong et al., 2017).

Chitinases are produced by many Paenibacillus spp. (Savita et al., 2013; Itoh et al., 2014; Ruth et al., 2016; Xiaoxiao et al., 2017 and Yong et al., 2017). Paenibacillus are known to have a biocontrol activity which due to their ability to produce variety of enzymes of degrading the plant cell wall, e.g. chitinases, cellulases, proteases and β-1,3-glucanases (Budi et al., 2000). Microbial chitinase are formulated as commercial biocontrol antifungal agents due to their ability to hydrolyze the cell walls of fungal pathogens (Flach et al., 1992 and Kim et al., 2008, 2010, 2011). Chitinase is part of the biological control agents (Chet et al., 1990 and Lorito et al., 1994) produced by isolates of Bacillus spp. (Chang et al., 2003; Lee et al., 2009 and Reyes-Ramirez et al., 2004), Paenibacillus spp. (Savita et al., 2013 and Itoh et al., 2014).

Although many studies mentioned isolation and characterization of Paenibacillus species with chitinolytic activity from different environments; hot spring soils (Hui et al., 2011), seafoods (Singh, 2010) and marine (Patel et al., 2007). A few numbersof studied focused on thermophilic bacteria having chitinolytic activities (Manucharova et al., 2011).

In recent years, thermostable chitinases received increased attention, because of their applicable properties in various industrial processes which performed at high temperatures (Kuzu et al., 2012). Soil microorganisms, especially thermophilic species have the ability to produce chitinolytic enzymes with thermostable nature (Manucharova et al., 2011).

Because of the various applications of chitinases, obtaining a novel chitinolytic bacterial strains with unique extreme properties is of great concern. Desert soils are harboring microorganisms growing at a broad temperature range. Therefore, the present study aims to isolate and characterize bacterial strains with thermostable chitinolytic activity from desert soils of Rafha governorate. Further, the physicochemical properties of the produced chitinase from this strain and its antifungal activity against some fungal hytopathogens were also determined.

**Materials and Methods**

**Materials**

Chitin (crab shell) and ρ-nitropheny-l-N-acetylglucosamine were purchased from Sigma-Aldrich Chemical Co. (St. Louis. MO, USA). All other chemicals used were of analytical grade purchased from either Sigma or Merck Chemical Co. (KGaA, Darmstadt, Germany).

**Sampling**

Ten soil samples were collected from the upper 5-15cm depth around wild plants; Horwoodia dicksoniae & Rumex cyprius from desert of Rafha governorate at the Northern Border region in Kingdom of Saudi Arabia. The collected samples were taken in clean plastic bags, until further proceedings, samples were stored in the refrigerator at 4°C.

**Preparation of colloidal chitin**

Colloidal chitin was prepared according to an established method of Chandrasekaran et al. (2012), 5g of chitin powder was added slowly to 100ml of concentrated HCl with vigorous stirring on ice overnight. Then, the mixture was added dropwise to 1L of ice-cold ethanol with rapid stirring for 4h. Finally, the precipitate was collected by suction filtration and washed with water until the washing solution was neutral. The colloidal chitin was stored at 4°C for subsequent use.

**Selective isolation of chitinase-producing bacteria**

For isolation of chitinase-producing bacterial...
strains, 5.0g soil were added to 100ml saline (0.85%) and shaken vigorously for 1h. Serial dilutions were made and spread onto colloidal chitin agar (CCA) media containing (w/v): 0.5% colloidal chitin, 0.7% (NH₄)₂SO₄, 0.1% K₂HPO₄, 0.1% NaCl, 0.01% MgSO₄·7H₂O, 0.05% yeast extract, 0.1% bacto tryptone and 2.0% agar at pH 7.0 (Yuli et al., 2004). The plates were incubated at 50°C for 7 days, and colonies showing clear halos were selected. Colonies exhibiting different morphological appearances were picked individually and further purified by subculturing on the same medium. Once purified, each isolate was processed to find out its chitin solubilizing (CS) ability in CC medium. This was done by calculating the zone of clearance to colony size (Faramarzi et al., 2009). Isolates were stored as a glycerol stock at -80°C. Among these strains, one designated as NBR10 exhibited high CS and was selected for further investigations.

**Taxonomic studies of chitinase-producing NBR10 strain**

**Morphological, physiological and biochemical characterization**

Morphological and physiological characterization of the isolate was performed according to the Bergey’s Manual of Systematic Bacteriology (Williams et al., 1989). Morphological characteristics such as colony morphology and cell morphology of the selected bacterial strain were studied (Holt et al., 1994). For biochemical characterization, different biochemical tests were studied by the test kit of api® 20E and api® 50 CHB Medium, bioMérieux, Marcy-l’Etoile, France. The results were analyzed with the APILAB Plus software (Logan & Berkeleym, 1984 and Fundagül et al., 2009). The test of api was carried out at Bacteriology Laboratory, the Holding Company for Biological Products and Vaccines (VACSERA), El-Dokki, Giza, Egypt. The biochemical tests were constructed according to methods described by Sneath (1984).

**Molecular characterization**

DNA isolation and amplification of 16S rRNA gene: A single, pure colony of the isolate was taken from the plate grown culture and was suspended in 10μl TE buffer [10mM Tris (pH 8.0), 1m MEDTA]. The samples were mixed briefly using cyclo mixer, and 5.0μl of this suspension was applied to FTA® Classic Card (Whatman International Ltd.) and allowed to dry for at least 1h at room temperature. Further processing of the sample was done according to the manufacturer’s instructions. The 16S rRNA genes were amplified using forward (5’-GAGTTTTGATCTGGCCTCAG-3’) and reverse (5’-AGAAAGGAGGTATCCAGCC’) primers (Xcelris Labs Ltd., India) known to be conserved among all known bacteria (Ulrike et al., 1989). PCR was performed in an automated thermal cycler (PTC-200, MJ Research Inc.) with an initial 92°C denaturation for 2min 10sec; followed by 35 cycles of 92°C for 1min 10sec, 48°C for 30sec, 72°C for 2min 10sec and a final extension at 72°C for 6min 10sec (Massol-Deya et al., 1995). The PCR products were resolved by electrophoresis on 1.5% agarose gel (BioRad, Hercules, CA) and visualized using gel documentation system (UVIpro, UVItec).DNA isolation and purification, 16S rRNA gene amplification and sequencing was carried out at Clinilab Company, El-Maadi, Cairo, Egypt.

**Phylogenetic analysis:** The partial sequences obtained from isolated strain was initially compared with reference sequences by using BLAST (National Centre for Biotechnology Information at www.ncbi.nlm.gov/BLAST) to determine their phylogenetic affiliations. The sequences were manually checked and submitted to Ribosomal Database Project (RDP at http://rdp.cme.msu.edu/) and the most closely related phylotype was identified. The sequences were finally aligned in the alignment explorer tool of the MEGA 4 software (Tamura et al., 2007) with the alignment parameters set to default values, and the phylogeny reconstruction analysis was performed with the help of the Neighbour joining method.

**Nucleotide sequence accession number:** The partial 16S rRNA gene sequence determined for chitinase-producing bacterial strain NBR10, identified as *Paenibacillus* sp., was deposited in the GenBank under accession number KT957624.1.

**In-vitro antifungal activity**

The antifungal activity of the strain NBR10 was investigated against three important fungal phytopathogens, *Alternaria burnsii*, *Fusarium oxysporum* and *Rhizoctonia solani*, under in vitro conditions using a dual culture technique (Faramarzi et al., 2009). These phytopathogens were supplied from Plant Protection Department, Desert Research Centre, Cairo, Egypt and maintained on potato dextrose agar (PDA, Merck Co.) medium.
The antifungal activity of the isolate NBR10 was determined on yeast malt extract agar (YMEA) medium by perpendicular streak method (Wadetwar & Patil, 2013). Isolated bacterial strain was cross streaked as a single line on solidified YMEA media in a petridish and incubated at 28°C for 72h. The fungal test organisms were then cross streaked perpendicular to the original streak of isolates. Control plates of the same medium without bacterial growth were also simultaneously streaked with test organism to study their normal growth.

**Chitinase production**

Chitinase production by the isolate NBR10 was performed at 50°C in 250ml Erlenmeyer flasks containing 50ml CC medium (pH 7.0) at 180rpm for 7 days, and the enzyme production was monitored at every 24h.

**Assay of chitinase activity**

The chitinase activity was determined colorimetrically by detecting the amount of N-acetyl-D-glucosamine (NAG) released from a colloidal chitin substrate (Vyas & Deshpande, 1989). A 48h grown culture was harvested and centrifuged in a microfuge (Biofuse Primo-R) at 10,000rpm for 10min to obtain cell-free culture supernatant. The reaction mixture (1.0ml) for enzyme assay consisted of enzyme (340μl) with CC (3mg) as a substrate in 50mM sodium acetate buffer (pH 5.0). This mixture was incubated at 55°C for 1h; the remaining colloidal chitin of the reaction was removed by centrifugation at 10,000rpm for 10min, and the chitinase activity was assayed in the supernatant by measuring the released sugar from colloidal chitin (Nelson, 1944). Briefly, the above supernatant was reacted with dinitrosalicylic acid (DNS) by boiling for 5min in absorbance of the released NAG was measured in reaction mixture at 540nm after cooling to room temperature. One international unit (IU) was defined as the activity that produced 1μmol of the product per hour.

**Physico-chemical characterization of enzyme**

Chitinase activity was measured in crude culture supernatant of isolate NBR10 at various pH values (3.0–10.0) using the following buffers (50mM): sodium acetate buffer (pH 3.0–6.0), sodium phosphate buffer (pH 7.0), Tris–HCl buffer (pH 8.0–9.0) and glycine-NaOH buffer (pH 10.0); temperature ranged from 20 to 100°C. The thermal stability was determined by initially preincubating enzyme at various temperatures (20–100°C) for different time intervals (0–44h) in a dry bath. After the heat treatment, samples were cooled and assayed for residual enzyme activity.

**Statistical analysis**

All the optimization studies were conducted in triplicate and the data were analyzed using single factor analysis of variance (ANOVA). All the data are graphically presented as the mean± S.D. of triplicates (n= 3). ANOVA was performed using Microsoft Excel 2007.

**Results and Discussion**

**Isolation and screening for chitinase producing bacteria**

Isolation of bacterial strains capable of producing chitinase with a novel property such as stability for a long time at high range of temperature, have been the focus of recent research. In this study, an attempt has been made to characterize thermos-stable chitinase enzyme produced by the soil-isolated bacterial strain. Eleven bacterial cultures were isolated from different soil samples collected from different localities from desert of Rafha governorate in Kingdom of Saudi Arabia. The obtained bacterial isolates were screened qualitatively for chitinase production under thermal (50°C) conditions using colloidal chitin agar (CCA) medium. Faramarzi et al. (2009) reported that, the qualitative assay of chitinase was performed by measuring the clear zone around the colony. Out of 11 isolates, 3 (27.2%) isolates were found to be chitinase positive through appearance of clear zone on the CC agar medium. Among them, strain NBR10 which isolated from soil sample from rhizosphere of wild plant, being the highest chitinase producer (Fig. 1). This isolate was selected for production of the enzyme in the fermentation medium. Soil rhizosphere contains many hydrolytic enzymes which included in recycling of nutrients, maintaining of soil fertility, decomposition of organic residues. The extracellular enzymes produced by rhizospheric microorganisms play initial role in degradation of high molecular weight polymers such as cellulose, pectin, lignin and chitin (Geetha et al., 2014).
Characterization of selected chitinolytic strain

Morphological, physiological and biochemical characterization

On the basis of morphology, isolate NBR10 showed morphological features typical for the genus *Paenibacillus*. The macroscopical investigation of colonies grown on nutrient agar plates showed smooth, translucent, convex and creamy coloured colonies. The microscopical examination showed small rod-shaped Gram-positive cells (Fig. 2).

The biochemical and physiological characteristics of this isolate NBR10 are recorded in Table 1. It was found that the isolate was able to grow at pH values between 5 and 10 with optimum pH 8 and in the presence of 2–6% of NaCl. The maximum growth temperature was recorded at 50°C, while the limiting growth was observed at 60°C. Also, the isolate was positive for amylase, caseinase, lipase and catalase enzymes. The results of biochemical and physiological characteristics of the isolate NBR10 recorded in Table 1 are analyzed with the APILAB Plus software. Our identification results are compared with data in Bergey’s Manual of Systematic Bacteriology (Sneath, 1984) and Bergey’s Manual of Determinative Bacteriology (Holt et al., 1994) of the genus *Paenibacillus*.

The morphological, biochemical and physiological identification of the isolate NBR10 was confirmed with sequencing of 16S rRNA which ensured relatedness of this isolate to the family Paenibacillaceae. The phylogenetic analysis of sequence of NBR10 was performed and phylogenetic tree is constructed (Fig. 3) with 98% similarity to *Paenibacillus thailandensis* S3-4A (NR_041490.1) hence our isolate is coded as *Paenibacillus* sp. NBR10. Ash et al. (1993) genus *Paenibacillus* includes facultative anaerobic, endospore-forming, Gram-positive bacterium previously was classified as *Bacillus* while in the recent time it classified as a separate genus.
**TABLE 1. Biochemical and physiological results of isolate NBR10 isolate.**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>NBR10</th>
<th>Characteristic</th>
<th>NBR10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>-</td>
<td>Glycogen</td>
<td>+</td>
</tr>
<tr>
<td>Starch</td>
<td>-</td>
<td>Erythritol</td>
<td>-</td>
</tr>
<tr>
<td>Gentio-bose</td>
<td>-</td>
<td>D-arabinose</td>
<td>+</td>
</tr>
<tr>
<td>Xylitol</td>
<td>-</td>
<td>Glycerol</td>
<td>-</td>
</tr>
<tr>
<td>D-lyxose</td>
<td>+</td>
<td>L-arabinose</td>
<td>+</td>
</tr>
<tr>
<td>D-turanose</td>
<td>-</td>
<td>D-xylose</td>
<td>+</td>
</tr>
<tr>
<td>D-fucose</td>
<td>-</td>
<td>D-ribose</td>
<td>+</td>
</tr>
<tr>
<td>D-tagatose</td>
<td>-</td>
<td>D-adonitol</td>
<td>-</td>
</tr>
<tr>
<td>D-arabitol</td>
<td>+</td>
<td>L-xylose</td>
<td>+</td>
</tr>
<tr>
<td>L-fucose</td>
<td>-</td>
<td>D-galactose</td>
<td>+</td>
</tr>
<tr>
<td>Gluconate</td>
<td>+</td>
<td>Methyl-β-D-Xylopyranoside</td>
<td>-</td>
</tr>
<tr>
<td>Fructose</td>
<td>+</td>
<td>L-arabitol</td>
<td>+</td>
</tr>
<tr>
<td>D-glucose</td>
<td>+</td>
<td>5-keto-gluconate</td>
<td>-</td>
</tr>
<tr>
<td>L-sorbose</td>
<td>-</td>
<td>2-keto-gluconate</td>
<td>-</td>
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<tr>
<td>D-mannose</td>
<td>-</td>
<td>Arginine dihydrolylase</td>
<td>-</td>
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<tr>
<td>D-lactitol</td>
<td>-</td>
<td>Beta-galactosidase</td>
<td>+</td>
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<tr>
<td>Ornithine decarboxylase</td>
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<td>L-rhamnose</td>
<td>+</td>
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<tr>
<td>Lysine decarboxylase</td>
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<td>D-mannitol</td>
<td>+</td>
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<tr>
<td>Hydrogen sulfide</td>
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<td>Inositol</td>
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<tr>
<td>Citrate</td>
<td>+</td>
<td>Methyl-β-D-Mannopyranoside</td>
<td>-</td>
</tr>
<tr>
<td>Tryptophane deaminase</td>
<td>+</td>
<td>D-sorbitol</td>
<td>-</td>
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<tr>
<td>Urease</td>
<td>+</td>
<td>Amygdalin</td>
<td>+</td>
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<tr>
<td>Indole</td>
<td>-</td>
<td>Methyl-β-D-Glucopyranoside</td>
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<tr>
<td>N-acetylglucosamine</td>
<td>-</td>
<td>Gelatinase</td>
<td>+</td>
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<tr>
<td>Esculin</td>
<td>+</td>
<td>VP</td>
<td>+</td>
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<tr>
<td>Arbutin</td>
<td>-</td>
<td>O-F test oxidation</td>
<td></td>
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<tr>
<td>D-cellobiose</td>
<td>+</td>
<td>Nitrate reduction</td>
<td>+</td>
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<tr>
<td>Salicin</td>
<td>+</td>
<td>Caseinase</td>
<td>+</td>
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<tr>
<td>D-lactose (bovine origin)</td>
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<td>Catalase</td>
<td>+</td>
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<tr>
<td>D-maltose</td>
<td>+</td>
<td>Lipase</td>
<td>+</td>
</tr>
<tr>
<td>D-saccharose (sucrose)</td>
<td>+</td>
<td>Nuclease</td>
<td>-</td>
</tr>
<tr>
<td>D-melibiose</td>
<td>+</td>
<td>Amylase</td>
<td>+</td>
</tr>
<tr>
<td>Inulin</td>
<td>-</td>
<td>Range of growth</td>
<td></td>
</tr>
<tr>
<td>D-trehalose</td>
<td>+</td>
<td>NaCl 2-6% (w/v)</td>
<td>+</td>
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<tr>
<td>D-raffinose</td>
<td>+</td>
<td>Temperature 20-60°C</td>
<td>+</td>
</tr>
<tr>
<td>D-melezitose</td>
<td>-</td>
<td>pH 7.10</td>
<td>+</td>
</tr>
</tbody>
</table>

+= Positive, - = Negative
Antifungal activity of Paenibacillus sp. NBR10

The local NBR10 isolate exhibited antifungal activity against 3 of phytopathogenic fungi with percent of growth inhibition arranged as 52.5% against *Fusarium oxysporum*, 75.0% against *Alternaria burnsi* and 85.71% against *Rhizoctonia solani* (Fig. 4) which indicate possibility of using of this strain as a biocontrol agent against fungal phytopathogens. *Paenibacillus* sp. BISR-047 exhibited similar growth inhibition against various phytopathogenic fungi: *F. oxysporum* and *A. burnsi* (Savita et al., 2013). Many species of *Paenibacillus* can be act as antagonists for plant pathogens, such as nematodes, bacteria and fungi (Bloemberg & Lugtenberg, 2001; Sheng et al., 2014 and Natalija & Dzoko, 2018).

Production of chitinase from Paenibacillus sp. NBR10

Impact of different incubation periods on production chitinase was investigated from 1 to 8 days at 50°C. The enzyme activity was recorded after 1st day of incubation, the maximum activity (2.71±0.076U/ml) was observed at 5th day (Fig. 5). Singh (2010) reported that, production of chitinase is proportional with concentration of chitin in the medium where after complete utilization of chitin results in decreasing in the chitinase activity. Many studies discussed the impact of incubation period on chitinase yield in different *Paenibacillus* species. Ruth et al. (2016) reported the incubation time required to reach the maximum enzyme level of chitinase production from the *Paenibacillus* sp. D3 was at 24h of incubation whereas, Singh (2010) reported the optimum incubation period for chitinase production from *Paenibacillus* sp. D1 was at 72h while it was 6 days for *Paenibacillus* sp. BISR-047 (Savita et al., 2013).

Effect of different temperatures on chitinase production

The incubation temperature incredibly influences microbial development rate, catalyst secretion, catalyst restraint, also protein denaturation (Kunamneni et al., 2005), the obtained chitinase showed high activity at a broad range of incubation temperature 20–60°C with optimum activity (2.91±0.152U/ml) at 55°C. A strong decrease in the chitinase activity (2.15±0.132U/ml) was recorded with increasing of temperature at 60°C (Fig. 6). Different studies focused on effect of temperature on chitinase production by *Paenibacillus* species and reported activity of these enzymes at different ranges. Singh (2010) reported activity of chitinase production by *Paenibacillus* sp. D1 at temperature range 30–60°C with optimum 50°C, whereas, Savita et al. (2013) reported that, chitinase production by *Paenibacillus* sp. BISR-047 was recorded at temperature range 28–65°C with optimum temperature 45°C.
Fig. 4. Antifungal activity of Paenibacillus sp. NBR10 against: (a) Alternaria burnii, (b) Rhizoctonia solani and (c) Fusarium oxysporum.

Fig. 5. Production of chitinase at different incubation period.

Fig. 6. Production of chitinase under different incubation temperatures.

In our study *Paenibacillus* sp. NBR10 which was found to grow at temperatures up to 60°C, but did not show any growth above 65°C, whereas its crude chitinase showed activity up to 55°C. This shows that the enzyme produced by thermophilic bacterial strain NBR10 is highly thermophilic in nature. The area selected for the present study has been characterized by complex dune physiography with alkaline soil pH and extremely low organic carbon content (Gothwal et al., 2007). In these areas, soil temperature crosses 50°C in the summers and this can be correlated with the growth temperature of the isolated strain.

**Effect of different pH values on chitinase production**

pH is one of the most important factors influencing the enzyme activity where the balance of the microbial metabolite depend on concentration of the hydrogen ion (Kunamneni et al., 2005). In this study, production of chitinase was recorded at pH range 3.0-8.0, with the maximum activity 2.63±0.104U/ml at pH 7.0 (Fig. 7). Several workers have reported broad range of pH optima for chitinase, from 4.0 to 10.0 for *Paenibacillus pasadenensis* CS0611 (Xiaoxiao et al., 2017), 4.0 to 10.0 for *Paenibacillus thermaeophilus* TC22-2b (Ueda & Kurosawa, 2015), from 3.0 to 9.0 for *Paenibacillus barengoltzii* (Fu et al., 2016).

**Effect of temperature and time on stability of chitinase**

Effect of thermal conditions on chitinase activity produced by *Paenibacillus* sp. NBR10 was studied through incubation of the enzyme in 50mM sodium acetate buffer (pH 7.0) at temperatures ranged from 20 to 100°C for 1h. The activity of the enzyme was determined at the range 30–90°C where the enzyme retains with more than 50% of its activity at this range. The activity of chitinase was increased gradually with increasing the temperature up to 60°C. The maximum activity was obtained when temperature increased to 65°C. But it showed slight decrease in its activity with the increasing in temperature at 70°C (Fig. 8). These results ensure the thermal nature of the produced enzyme.

The produced chitinase was found to be stable when incubated for period of time ranged from 0 to 36h in pH 7.0 and 65°C, where no change in chitinase activity was observed when the enzyme incubated in these conditions (Fig. 9). Although the enzyme activity decreased when incubation time increased for more than 36h, 59% of original enzyme activity was retained. The obtained results ensured stability of chitinase enzyme produced by *Paenibacillus* sp. NBR10 in thermal conditions for a long period of time. Similar results have been reported for many works of *Paenibacillus* species such as *Paenibacillus* sp. (Singh, 2010), *Paenibacillus* sp. BISR-047 (Savita et al., 2013), *Paenibacillus barengoltzii* (Fu et al., 2016).

Most of the industrial processes which depend on enzymatic reactions are performed at high temperatures, thus in the last few years, researches focusing on thermostable enzymes has been increased. As we know, there is no previous reports on isolation and characterization of thermostable chitinase by *Paenibacillus* sp. from the Northern Border region, in Saudi Arabia.
Conclusion

Conventionally, fungicides used to control damage caused by fungal phytopathogens contribute several environmental problems as degradation, pollution, lethal effect on beneficial rhizobacteria and development of resistant strains. Thus, there is a great need for alternate strategies for controlling the fungal diseases. Thermotolerant chitinase-producing bacteria offers a potential alternate and additive to toxic chemical fungicides in reducing their dosage in agriculture. Our results conclude, the locally isolated *Paenibacillus* sp. NBR10 isolate exhibited unique properties of potent inhibition of phytopathogenic fungi and high chitinase production. Hence to confirm these properties especially under field conditions further *in vivo* studies are required.

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References


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فعالية بكتيريا بينيباسيلليس نوع ان بي آر10 المحبة للحرارة والمعزولة من التربة كعامل تحكم حيوي مخلل للكيتيين

محمد السيد(1)، ألقت نصار(1)، عبد الناصر قبيصى(1)

أقسم الأحياء – كلية العلوم والآداب – جامعة الحدود الشمالية – المملكة العربية السعودية،(2) قسم النبات
الكيميائي – كلية العلوم – جامعة الأزهر – القاهرة – مصر،(3) قسم الاقتصاد المنزلي (تقنية) – كلية
العلوم والآداب – جامعة الحدود الشمالية – المملكة العربية السعودية،(4) قسم وقاية النبات – مركز بحوث

حظيت إنزيمات الكيتيناز الميكروبية باهتمام متزايد بسبب نطاقها الواسع من تطبيقات التقنية الحيوية خاصة
في الزراعة من أجل المكافحة الحيوية للفطريات الممرضة للنبات. تنتج إنزيمات الكايتينز بواسطة عدد من
الكائنات الدقيقة، بما في ذلك البكتيريا. وبسبب الاهتمام المتزايد في إنزيمات الكايتينز الثابت حراريا، هناك حاجة
مستمرة لعزل وتوصيف سلالات بكتيرية جديدة لها خصائص فريدة من مثل هذه الأنزيمات. في هذه الدراسة، تم
وصف وتعريف ودراسة الخصائص الفيزيوكيميائية لأنزيم الكايتينز الذي تنتج السمالة البكتيرية المحبة للحرارة
إن بي آر10. والمعزولة من التربة الصحراوية في محافظة رفحاء في منطقة الحدود الشمالية في المملكة العربية
السعودية. تم تعريف السمالة المنتجة لألزم الكايتينز نوع من بينيباسيلليس ان بي آر10. على أساس الخصائص
المورفولوجية والكيميائية والكيميائية الحيوية بالإضافة إلى تحليل التتابع النيوكليتيدي التسلسلي لجين الحمض
الرئوي الريبوسومي 16 اس ( الرقم الدخول في بنك الجينات KT957624.1) وجد أن السلاله المعزولة لها
نشاط قوي مضاد لثلاثة من الفطريات الممرضة للنبات الالترناريا بورسني، رايزكتونيا سولاني و
على التوالي. إنزيم الكايتينز الذي تم الحصول عليه
من نوع بينيباسيلليس ان بي آر10. أظهر نشاطاً على نطاق واسع من الأس الهيدروجيني (4-9) ودرجات
حرارة (30-90 درجة مئوية)، مع نشاط أقصى عند 7.0 درجة محتوية درجة حرارة 55 درجة مئوية
على التوالي. أيضًا، وجد أن الأنزيم المنتج ثابت حراريًا عند درجات حرارة أعلى، حيث يحتفظ بـ 90% من نشاطه
عند 65 درجة مئوية لمدة 36 ساعة التنشيط الكيتوتستريغ والمواد للطريقة من العزلة المحلية بينيباسيلليس ان
بي آر10. ترشح هذه السلاله لاستخدامها كعامل تحكم بيولوجيًا فعالًا.